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# Programmed Cell Death in *Neurospora crassa* Is Controlled by the Allorecognition Determinant *rcd-1*

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**ABSTRACT** Nonself recognition following cell fusion between genetically distinct individuals of the same species in filamentous fungi often results in a programmed cell death (PCD) reaction, where the heterokaryotic fusion cell is compartmentalized and rapidly killed. The allorecognition process plays a key role as a defense mechanism that restricts genome exploitation, resource plundering, and the spread of deleterious senescence plasmids and mycoviruses. Although a number of incompatibility systems have been described that function in mature hyphae, less is known about the PCD pathways in asexual spores, which represent the main infectious unit in various human and plant fungal pathogens. Here, we report the identification of *regulator of cell death-1* (*rcd-1*), a novel allorecognition gene, controlling PCD in germinating asexual spores of *Neurospora crassa*; *rcd-1* is one of the most polymorphic genes in the genomes of wild *N. crassa* isolates. The coexpression of two antagonistic *rcd-1-1* and *rcd-1-2* alleles was necessary and sufficient to trigger cell death in fused germlings and in hyphae. Based on analysis of wild populations of *N. crassa* and *N. discreta*, *rcd-1* alleles appeared to be under balancing selection and associated with *trans*-species polymorphisms. We shed light on genomic rearrangements that could have led to the emergence of the incompatibility system in *Neurospora* and show that *rcd-1* belongs to a much larger gene family in fungi. Overall, our work contributes toward a better understanding of allorecognition and PCD in an underexplored developmental stage of filamentous fungi.

**KEYWORDS** *Neurospora*; programmed cell death; allorecognition; heterokaryon incompatibility; cell fusion

**A**LLORECOGNITION or discrimination against conspecific but genetically distinct nonself is a phenomenon that is important in various colonial organisms that limits genome exploitation and resource plundering (Debets and Griffiths 1998; Laird *et al.* 2005; Ho *et al.* 2013; Bastiaans *et al.* 2016). The phenomenon has been described in early diverging invertebrate metazoans, including the ascidian *Botryllus schlosseri*, the cnidarian *Hydractinia symbiolongicarpus*, and in the social amoebae *Dictyostelium discoideum*, where it restricts

colonial chimerism to closely related individuals (Laird *et al.* 2005; Lakkis *et al.* 2008; Rosengarten and Nicotra 2011). In filamentous fungi, where cell fusion (anastomosis) between genetically different fungal colonies is often abortive, allorecognition mechanisms play a similar role by triggering a programmed cell death (PCD) reaction termed vegetative or heterokaryon incompatibility (HI) (Glass and Dementhon 2006; Paoletti 2016; Daskalov *et al.* 2017). The cell death reaction is spatially and temporally restricted to the heterokaryotic fusion cells in the contact zone between two genetically distinct individuals, and is often associated with the formation of a pigmented demarcation line or barrage between incompatible strains (Perkins *et al.* 2001). Fusion cells undergo extensive vacuolization, hyper-septation, ROS production, lipid droplet accumulation, and plasma membrane disruption (Saupe *et al.* 2000; Glass and Kaneko 2003). HI has been described in dozens of filamentous ascomycete species, and the populations of numerous phytopathogenic species have been characterized in terms of vegetative compatibility groups

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(VCGs). Strains belonging to the same VCG are able to form heterokaryons and transmit genetic content horizontally and often exhibit similar virulence and host specificity, while strains of different VCGs are incompatible and do not form viable heterokaryons (Zeise and Von Tiedemann 2002; Fan *et al.* 2018). By restricting chimerism between individuals, HI limits the horizontal transmission of deleterious cytoplasmic elements (*i.e.*, mycoviruses, senescence plasmids), thus functioning as a fungal defense mechanism (Debets *et al.* 1994; Zhang *et al.* 2014). In *Cryphonectria parasitica*, the causal agent of chestnut blight disease, systematic disruption of allorecognition genes that prevented heterokaryon formation between strains of different VCG resulted in the creation of a super mycovirus donor strain (Zhang and Nuss 2016). The “super mycovirus donor strain” carries a hypo-virulence plasmid, and, hence, can be an effective biocontrol agent (Zhang and Nuss 2016). Similar approaches underscore the importance of genetically identifying and characterizing novel allorecognition loci in fungi.

*Neurospora crassa* is a model filamentous ascomycete species for studying cell–cell communication, anastomosis formation, and HI (Herzog *et al.* 2015; Zhao *et al.* 2015; Daskalov *et al.* 2017). At least 11 loci control HI in *N. crassa* (Mylyk 1975). Several *het* genes (for HI) have been molecularly identified and shown to induce cell death when coexpressed with incompatible alleles of the same gene (allelic HI systems) or with incompatible alleles of a different gene (nonallelic HI systems) (Kaneko *et al.* 2006; Paoletti 2016). The identified *het* genes are often highly polymorphic, with several different alleles segregating in near equal frequencies in wild populations of *N. crassa* (Wu *et al.* 1998; Hall *et al.* 2010). Such a distribution pattern of antagonistic alleles in wild isolates is an indication that a particular locus is under balancing selection. Balancing selection has been described to operate on the self-incompatibility locus (SI) in flowering plants, the major histocompatibility complex (MHC) in mammals, and has been associated with allorecognition loci in other animals (Richman 2000; Nydam *et al.* 2017). Alleles of a gene under balancing selection can be maintained in populations undergoing speciation events, which results in a *trans*-species segregation of multiple allelic haplogroups (Klein *et al.* 1998). Evolutionary hallmarks of long-term balancing selection and *trans*-species polymorphism have been recently used in a population genomics analysis of *N. crassa* wild isolates to successfully identify novel *het* genes (Zhao *et al.* 2015). However, in spite of the frequently shared molecular hallmarks of evolution, *het* gene function seems poorly conserved between species (Paoletti 2016).

The products of several *het* genes belong to a large family of fungal proteins, analogous to NOD-like receptors (NLRs) (Dyrka *et al.* 2014). NLR proteins (also known as nucleotide-binding site and leucine-rich repeats containing proteins or NBs-LRR) are intracellular innate immune receptors controlling cell death in plants and animals in response to immunogenic cues (Duxbury *et al.* 2016; Jones *et al.* 2016). The similarities between *het* determinants and genes involved in innate immunity in other eukaryotic lineages have prompted

the hypothesis that HI may have originated from signaling pathways mediating interspecific biotic interactions, possibly in the context of an uncharacterized fungal innate immunity system (Paoletti and Saupe 2009; Dyrka *et al.* 2014; Uehling *et al.* 2017). Thus, the molecular identification and characterization of novel allorecognition factors in fungi could help us elucidate the evolutionary history of nonself recognition processes in Eukaryotes.

Previously identified HI-inducing systems are suppressed during the *Neurospora* sexual cycle, and also in germinating conidia (germlings), which play a fundamental role for vegetative propagation (Shiu and Glass 1999; Ishikawa *et al.* 2012). However, recently Heller *et al.* (2018) identified a gene pair controlling PCD and allorecognition in germlings of *N. crassa*. Germling-regulated death (GRD) is triggered when germlings of incompatible haplotypes undergo cellular fusions during early colony establishment. The cell death reaction occurs very rapidly (~20 min) with the appearance of large vacuoles in the cytoplasm of fused incompatible germlings preceding cellular lysis and uptake of vital dyes (Heller *et al.* 2018). The two genes controlling GRD are linked in the genome of *Neurospora*. One gene (*plp-1*) encodes an NLR-like protein containing a phospholipase domain, and the other (*sec-9*) encodes a homolog of *SEC9* from yeast, a protein involved in plasma membrane fusion and exocytosis (Brennwald *et al.* 1994; Heller *et al.* 2018).

The discovery of GRD represents an exciting possibility to study fungal PCD in a well-controlled biological system and during a developmental stage of the fungal life cycle (asexual spores) of great importance for clinical and agricultural applications, because it constitutes the infectious stage of many human and plant fungal pathogens (Brown *et al.* 2012). Here, we further investigated the molecular basis of GRD, and identified a novel genetic determinant of allorecognition in *N. crassa*. By using bulked segregant analysis (BSA) of GRD-specific DNA pools and population genomics data, the NCU05712 locus was identified as a second GRD-controlling allorecognition factor, termed *regulator of cell death-1* (*rcd-1*). Viable germling fusion of cells containing incompatible alleles of *rcd-1* (*rcd-1-1* and *rcd-1-2*) was prevented by PCD induction. The *rcd-1* alleles are highly polymorphic in *N. crassa* populations and show signs of balancing selection. Furthermore, we identified genomic rearrangements in the *rcd-1* locus, which could be at the origin of this allorecognition system in *Neurospora*. Such genomic rearrangements could represent a general evolutionary mode for fungal incompatibility systems. Moreover, we show that *rcd-1* belongs to a large gene family in filamentous ascomycete species. The latter observation suggests that *rcd-1* might be an important component of cell death pathways in fungi with broader role in nonself recognition.

## Materials and Methods

### Strains, growth media, and molecular constructs

Strains were grown using standard procedures and protocols that can be found on the *Neurospora* homepage at FGSC (<http://www.fgsc.net/Neurospora/NeurosporaProtocolGuide.htm>).

Vogel's minimal media (VMM) (with supplements, if required) was used to culture all strains, except when specified otherwise (Vogel 1956). Crosses were performed on Westergaard's synthetic cross medium (Westergaard and Mitchell 1947). For flow cytometry experiments, thermo-reversible solid Vogel's medium was obtained substituting the agar with 20% Pluronic F-127 (Sigma-Aldrich).

All wild *N. crassa* strains in the present study were isolated from Louisiana, have been described previously, and are available at the FGSC (Dettman *et al.* 2003; Palma-Guerrero *et al.* 2013; Zhao *et al.* 2015; Corcoran *et al.* 2016) (Supplemental Material, Table S1). All engineered strains were from the genetic background of the wild-type laboratory strain (FGSC2489; OR74A). The  $\Delta$ NCU05712 deletion strain was obtained from the single gene deletion collection of *N. crassa* strains at the FGSC (Colot *et al.* 2006).

The *rcd-1-1* and *rcd-1-2* alleles with native promoters were cloned in the pMF272 vector (Freitag *et al.* 2004) using the restriction enzymes *NotI* and *ApaI* and introduced in the *his-3* locus of a  $\Delta$ *rcd-1* strain. Molecular fusions of *rcd-1* with fluorescent proteins (GFP or mCherry) were produced by cloning the *rcd-1* alleles in pMF272-derived vector using *MscI*/*PacI* (for the *rcd-1-1* allele) and *XbaI*/*PacI* (for the *rcd-1-2* allele) under the regulation of the *tef-1* promoter. Positive transformants were backcrossed with an  $\Delta$ *rcd-1* strain to construct homokaryotic strains that were subsequently verified by PCR.

### Bulked segregant analyses and genome resequencing

Bulked segregant analysis was performed as described (Heller *et al.* 2018). The GRD phenotype of progeny from a cross between FGSC2489 and a wild isolate JW258 (FGSC 10679) was determined by pairing germings with each parental strain and assessing PCD. Two GRD-specific DNA pools were produced (60 ng from 50 progeny strains in each DNA pool) for library preparation and sequencing. All paired-end libraries were sequenced on a HiSeq2000 sequencing platform using standard Illumina operating procedures (Vincent J. Coates Genomics Sequencing Laboratory, University of California, Berkeley). The mapped reads for each group of 50 pooled segregants are available at the Sequence Read Archive (PRJNA556444; <https://www.ncbi.nlm.nih.gov/sra/PRJNA556444>).

### Flow cytometry

Flow cytometry was performed according to Heller *et al.* (2018). We recorded 20,000 events per sample for each experiment. Experiments were performed at least three times. Data were analyzed with custom MATLAB (MathWorks) script with ungerminated conidia excluded from the analyses (Gonçalves *et al.* 2019). Cell death is shown as the average percentage of fluorescent events from all experiments.

### Evolutionary analysis of DNA and protein sequences

We characterized homologs of NCU05712 in a collection of 194 publicly available Sordariales genomes representing 16 - *Neurospora* species (Lasiosphaeriaceae II) (Kruys *et al.* 2015),

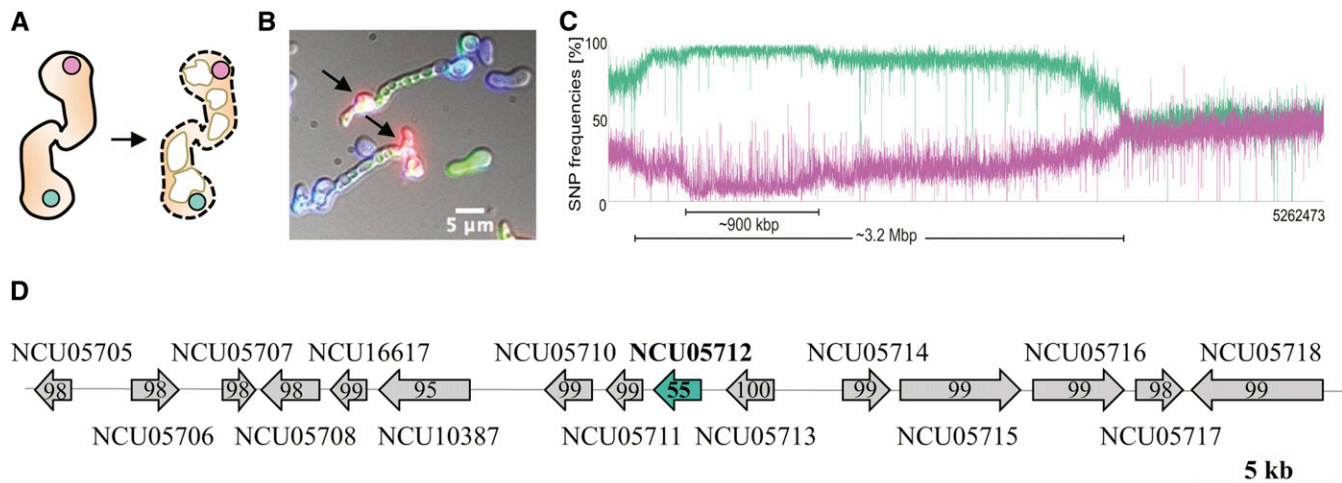
one non-*Neurospora* Lasiosphaeriaceae II species, two species in Lasiosphaeriaceae IV, one species in Lasiosphaeriaceae I, and 14 species of thermophilic subfamily Chaetomiaceae (Table S1). The dataset included resequencing data for 27 *N. crassa* isolates from Louisiana (Galagan *et al.* 2003; Zhao *et al.* 2015); 92 *N. tetrasperma* strains from North America, Europe, and Oceania (Ellison *et al.* 2011; Corcoran *et al.* 2016); and 43 *N. discreta* PS4 from North America, Europe, and Asia (Gladieux *et al.* 2015). We used publicly available genome assemblies and predicted genes where possible. Other genomes were assembled using ABySS (Simpson *et al.* 2009), and annotated using Augustus, with *N. crassa* as the reference species and intron locations obtained by mapping other species proteins onto each focal genome using Exonerate (Slater and Birney 2005).

We identified NCU05712 homologs in *Neurospora* genomes using BlastP against predicted proteins and keeping hits with an e-value <1e-5 and score >300. The resulting set of protein sequences was filtered from proteins whose length was >270 aa, aligned using Clustalo (Sievers and Higgins 2018), and used to build a hidden Markov model (HMM) profile using hmmbuild (Finn *et al.* 2011). We then searched for homologs in non-*Neurospora* genomes using hmmsearch (Finn *et al.* 2011), keeping only proteins in the inclusion threshold (Table S2). We removed from the dataset homologs identified in *Neurospora tetrasperma* reference genome, because gene models predicted premature stops codons. *N. tetrasperma rcd-1* ORF sequences were aligned using BLAST Global align (at NCBI), and sequences with SNPs or micro indels (up to six nucleotides) leading to predicted stop codons in the *N. tetrasperma rcd-1* ORFs were removed.

For each taxon with resequencing data, genomic variation at *rcd-1* was compared to a set of orthologous genes. Orthologous relationships were analyzed using Orthofinder (Emms and Kelly 2015) using Diamond (Buchfink *et al.* 2015) as the sequence aligner. We generated sets of single-copy orthologs present in 95% of isolates, and removed coding sequences not starting with "ATG" or having >10 Ns. TranslatorX (Abascal *et al.* 2010) and aligned coding sequences, while keeping the coding frame (protein sequence aligner: Muscle; sequence cleaning: GBlocks with parameters -g "-b2 8 -b3 3 -b4 3 -b5 n"). The number of protein variants was estimated based on nonsynonymous substitutions using Biopython, Ksmax (the maximum number of synonymous substitutions between sequences pairs) using yn00 in PAML (Yang 2007), and  $\pi$  (the average number of nucleotide differences between sequence pairs), s (number of segregating sites standardized by sequence length), and Tajima's D (a measure of skewness of the allele frequency spectrum) using Egglib v3 (De Mita and Siol 2012). Gene genealogy of *rcd-1* was inferred using the GTRGAMMA model in RAxML v8.2.10 (Stamatakis 2014). The genome genealogy within Sordariaceae used the same approach and program by concatenating coding sequences at 874 single copy orthologs present in at least 90% of isolates.

Long-term positive selection was assessed using a maximum likelihood framework for parameter inference and





**Figure 1** Bulk segregant analysis (BSA) and comparative genomics identifies GRD-controlling candidate gene. (A) Cartoon of germling-regulated death (GRD). Shown are two genetically incompatible germlings, represented by nuclei (circles) in pink and green, undergoing cellular fusion (left). The GRD reaction (right) manifests with extensive vacuolization of the germlings (white shapes) and cell lysis (shown with dashed line). (B) GRD reaction occurring between genetically incompatible asexual spores of two different strains as revealed by the uptake of the vital dye propidium iodide (PI, black arrows) at fusion points between incompatible germlings. One strain is marked with calcofluor-white (blue) and the other strain expresses cytoplasmic GFP (green). Bar, 5  $\mu$ m. (C) SNP segregation on linkage group III after sequencing of two genomic DNA pools of 50 segregant strains each from the FGSC2489 X JW258 cross. Purple line: SNP frequencies of pooled DNA of segregant strains showing GRD with FGSC2489. Green line: SNP frequencies of pooled DNA of segregant strains compatible with FGSC2489. (D) Genomic organization of the locus containing the GRD candidate gene NCU05712 or *regulator of cell death-1*. Percentage of identity between the parental strains (FGSC2489 and JW258) is shown for each ORF in the region.

codon-based substitution models that allow the dN/dS ratio  $\omega$  to vary among sites and among branches, as implemented in PAMs codeml program. For the site-specific analysis, we compared three codon substitution models: M7 which is a model of purifying selection that allows individual sites to evolve under differing levels of constraint and assumes a beta distribution of negatively selected ( $0 < \omega < 1$ ), M8a which is a model of purifying selection that adds neutral sites  $\omega = 1$  to M7, and M8, which is a model of positive selection that adds positively selected sites ( $\omega > 1$ ) to M7. For the branch-site analysis, assuming variable selective pressures among branches and sites, we specified *rcd-1-2* as the foreground branch and *rcd-1-1* as the background branch. We compared model A1, which is a model of purifying selection ( $0 < \omega \leq 1$ ) with model A, which adds positively selected sites along the foreground branch ( $\omega > 1$ ) to model A1. Nested models were compared using likelihood ratio tests.

To gather *rcd-1* homologs outside the Sordariales, we used the HMMER web server (Finn *et al.* 2011). The initial query sequence was *rcd-1-1* from *N. crassa* strain FGSC2489. Three iterative searches were performed to retrieve as many *rcd-1* homologs as possible (Table S2). Significant hits were aligned with Clustal, sorted by size, and sequences  $<230$  aa and  $>350$  were discarded. Sequences were assigned to taxonomic groups (Order) using NCBI Taxonomy, followed by manual curation of “Unclassified” taxa using Index Fungorum. *Pseudogymnoascus destructans* accounted for most of the sequences left “Unclassified” after manual curation. Protein sequences were aligned using Clustal, and a protein genealogy was inferred using model PROTGAMMAJTT in RAxML v8.2.10.

## Microscopy

Microscopy was performed on Zeiss Axioskop two equipped with a Q Imaging Retiga-2000R camera (Surrey), using a 40 $\times$ /1.30 Plan-Neofluar oil immersion objective and the iVision Mac4.5 software. Agar squares ( $\sim 1$  cm<sup>2</sup>) were excised from plates with growing germlings and observed under the microscope after 4–5 hr of growth (Heller *et al.* 2016).

## Data availability

The authors state that all data necessary for confirming the conclusions presented in the article are represented fully within the article. Mapped sequencing reads for BSA analysis of pooled segregant strains are available at the Sequence Read Archive (PRJNA556444). Files used for evolutionary analysis of *rcd-1* homologs are deposited at figshare and are publicly accessible with the following link <https://doi.org/10.6084/m9.figshare.10298045>. The deposited datasets are as follows – File 1: DNA sequences of *rcd-1* homologs in Sordariaceae used for phylogeny in Figure 5 and positive selection analyses. File 2: Protein sequences of *rcd-1* homologs in fungi and bacteria used for phylogeny presented in Figure 6. File 3: DNA sequences of genes representing the genomic background in *N. crassa* used for analyses of balancing selection at *rcd-1* in *N. crassa*. File 4: DNA sequences of genes representing the genomic background in *N. discreta* used for analyses of balancing selection at *rcd-1* in *N. discreta*. File 5: DNA sequences of genes representing the genomic background in *N. tetrasperma* used for analyses of balancing selection at *rcd-1* in *N. tetrasperma*. Raw flow cytometry data files are available upon demand. Supplementary Tables and

**Table 1 Average summary statistics (and 95% percentile) for *rcd-1* and reference genes representing the genomic background in three *Neurospora* species**

Species/gene set	$S^a$	$\pi^b$	$D^c$	$K_Smax^d$	$Pvar^e$
<i>Neurospora crassa</i>					
<i>rcd-1</i>	0.451**	0.220***	3.491***	1.762**	22.0***
Reference genes ( $n = 5632$ )	0.043 (0.136)	0.011 (0.031)	−0.490 (1.072)	0.261 (0.248)	8.0 (18.0)
<i>Neurospora discreta</i>					
<i>rcd-1</i>	0.491**	0.202***	2.637**	2.318**	16.0**
Reference genes ( $n = 5265$ )	0.032 (0.085)	0.008 (0.019)	0.244 (1.580)	0.324 (0.149)	5.7 (12.0)
<i>Neurospora tetrasperma</i>					
<i>rcd-1</i>	0.116	0.023	−0.908	0.218	6.0
Reference genes ( $n = 4698$ )	0.054 (0.154)	0.011 (0.032)	−0.0367 (2.190)	0.480 (0.319)	7.735 (17.0)

<sup>a</sup> Number of polymorphic sites per base pair.

<sup>b</sup> Average number of nucleotide differences between sequence pairs.

<sup>c</sup> Tajima's neutrality statistic (Tajima 1989).

<sup>d</sup> Maximum number of synonymous substitutions between sequences pairs.

<sup>e</sup> Number of amino-acid sequences based on nonsynonymous substitutions (\*\*\* in 99.9% percentile, \*\* in 99% percentile, \* in 95% percentile).

Figures are available at figshare: <https://doi.org/10.25386/genetics.10162688>.

## Results

### **BSA and population genomics identify NCU05712 as a highly polymorphic candidate gene controlling GRD**

To identify novel GRD-inducing genes, we used progeny from a cross between two wild isolates, the laboratory reference strain FGSC2489 and JW258 (FGSC10679). Genomic analyses of the progeny from this cross (FGSC2489 x JW258) were used previously to identify the kind of recognition determinants that act during chemotropic interactions between germlings and two linked GRD-inducing genes, *plp-1* and *sec-9* (Heller *et al.* 2016, 2018). The GRD phenotype (Figure 1, A and B) segregated 3:1 in progeny of this cross, suggesting that two independent loci regulated the GRD phenotype (*sec-9/plp-1* and a second unidentified locus). To identify the second GRD locus, we performed a BSA with progeny from an F2 backcross between FGSC2489 and an F1 progeny (Segregant 121) that showed GRD with FGSC2489 in spite of being isogenic to FGSC2489 at *sec-9/plp-1* (GRD due to second locus). More than 100 F2 progeny strains were phenotyped for GRD in cocultures with germlings of FGSC2489. Approximately half of the progeny formed viable heterokaryons with FGSC2489 germlings, while the other half showed rapid death of germlings ~20 min postfusion (Video 1). We extracted DNA from F2 progeny strains and sequenced two GRD-specific DNA pools (compatible with FGSC2489 and incompatible with FGSC2489) for BSA analysis. A random SNP distribution of ~50% was observed for the two groups on all chromosomes with the exception of a large region (~900 kbp) on chromosome III, where SNPs segregated at ~100% frequency between GRD groups (Figure 1C). Within this interval, we searched for highly polymorphic genes, hypothesizing that strongly divergent alleles might represent good candidates for being involved in allorecognition. While most genes in this region showed

high identity (>95%) between the two parental strains (FGSC2489 and JW258), NCU05712 was highly polymorphic and parental alleles showed only 55% identity in the ORF region (Figure 1D). The high degree of sequence divergence between the two allelic haplogroups of NCU05712 was not observed at any of the adjacent genes, which showed identity percentage ranging from 95 to 100%.

We asked how the observed level of polymorphism in NCU05712 compared to a set of reference genes representing the broader gene polymorphism found in the genomes of *N. crassa* isolates from Louisiana (the population of origin of the laboratory reference strain FGSC2489). We computed five summary statistics of genetic variation for the set of reference genes, and compared the scores obtained for each statistic, representing the overall polymorphism level in the genome, to scores obtained with NCU05712 for the same five statistics (Table 1). Our analysis ranked the GRD candidate gene NCU05712 in the top 1% of most polymorphic genes in the *N. crassa* genome for all five statistics (Table 1). Alleles at NCU05712 were in the 99.9% percentile on three of the computed five statistics—the number of amino acid sequences based on nonsynonymous substitutions (*Pvar*), the average number of nucleotide difference between sequences ( $\pi$ ), and Tajima's neutrality statistic *D* (Table 1). NCU05712 featured equally among the most polymorphic genes in *Neurospora discreta*, but not in *Neurospora tetrasperma*, consistent with the fact that we could retrieve only one of the NCU05712 allelic lines in the latter species (Table 1). These results indicate that NCU05712 is one of the most polymorphic genes in the genomes of at least two *Neurospora* species. Based on our analysis, we hypothesized that NCU05712 was the likeliest candidate gene to control GRD in the segregating region.

### **NCU05712 is necessary and sufficient for GRD**

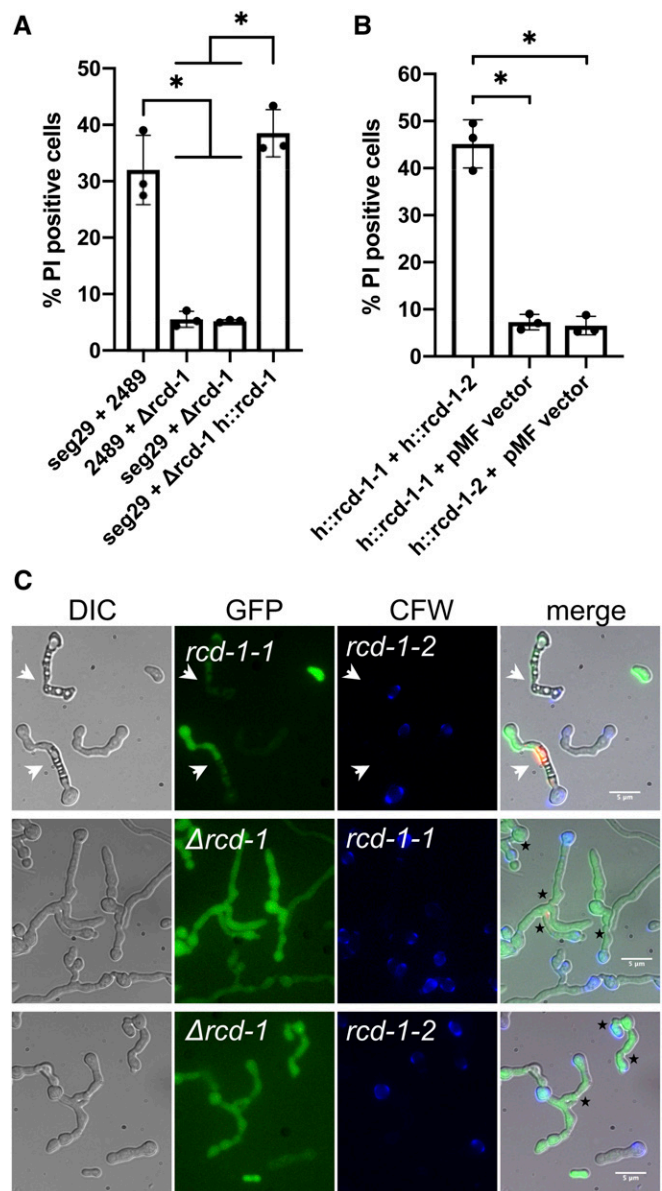
To test the hypothesis that NCU05712 controls GRD, a NCU05712 deletion strain from the *Neurospora* deletion collection (Colot *et al.* 2006; Dunlap *et al.* 2007) was characterized. First, we showed that the ΔNCU05712 mutant was not

affected in growth or conidiation as compared to the parental strain (FGSC2489) (Figure S1). We then compared the frequency of GRD in pairings between FGSC2489 germlings with an incompatible segregant strain (seg29) carrying the JW258 allele of NCU05712 or in pairs of  $\Delta$ NCU05712 germlings with seg29 using flow cytometry and the vital dye propidium iodide (PI) (Figure 2A; Video 1). When germlings of FGSC2489 were mixed in 1:1 ratio with germlings from seg29, we observed ~30% of pairs that exhibited PI uptake (a proxy for GRD). This frequency of GRD was significantly higher than that observed for mixtures of  $\Delta$ NCU05712 and seg29 germlings, where GRD was ~5% (Figure 2A); ~5% cell death was also observed in germlings undergoing self-fusion in a single strain (Figure S2). The strong reduction in GRD in  $\Delta$ NCU05712 + seg29 pairings segregated with the hygromycin resistance marker used to construct the deletion strain (Figure S3). Furthermore, the GRD phenotype was restored when NCU05712 was reintroduced into the  $\Delta$ NCU05712 strain (Figure 2A). We concluded that NCU05712 was necessary for the allorecognition reaction, and named the locus *regulator of cell death* or *rcd-1*. We termed the two allelic haplogroups as *rcd-1-1* (as represented by FGSC2489) and *rcd-1-2* (as represented by JW258).

To test if polymorphism at the *rcd-1* locus was sufficient to trigger cell death, we targeted an *rcd-1-1* allele or an *rcd-1-2* allele to the *his-3* locus of a  $\Delta$ *rcd-1* strain. Paired germlings of engineered *rcd-1-1* and *rcd-1-2* isogenic strains, expressing the antagonistic *rcd-1* alleles in otherwise isogenic backgrounds, produced a strong GRD reaction with a ~45% cell death frequency (Figure 2B). Strains expressing just *rcd-1-1* or *rcd-1-2* did not show GRD above the background level when grown by themselves or in cocultivations with a  $\Delta$ *rcd-1* strain transformed by an empty control vector (pMF274) (Figure 2, B and C and Figure S3). In addition, we observed an incompatible reaction in ascospore progeny bearing both *rcd-1-1* and *rcd-1-2* (Figure S4), while heterokaryotic fusion cells bearing fluorescently tagged RCD-1-1 and RCD-1-2 showed strong vacuolization and compartmentation of fused cells. These results indicated that allelic differences at the *rcd-1* locus were necessary and sufficient to trigger cell death in fused germling pairs and also in hyphae in *N. crassa*, and established *rcd-1* as a second allorecognition system controlling GRD in *N. crassa*.

#### ***rcd-1* does not act downstream of *plp-1* in the induction of GRD**

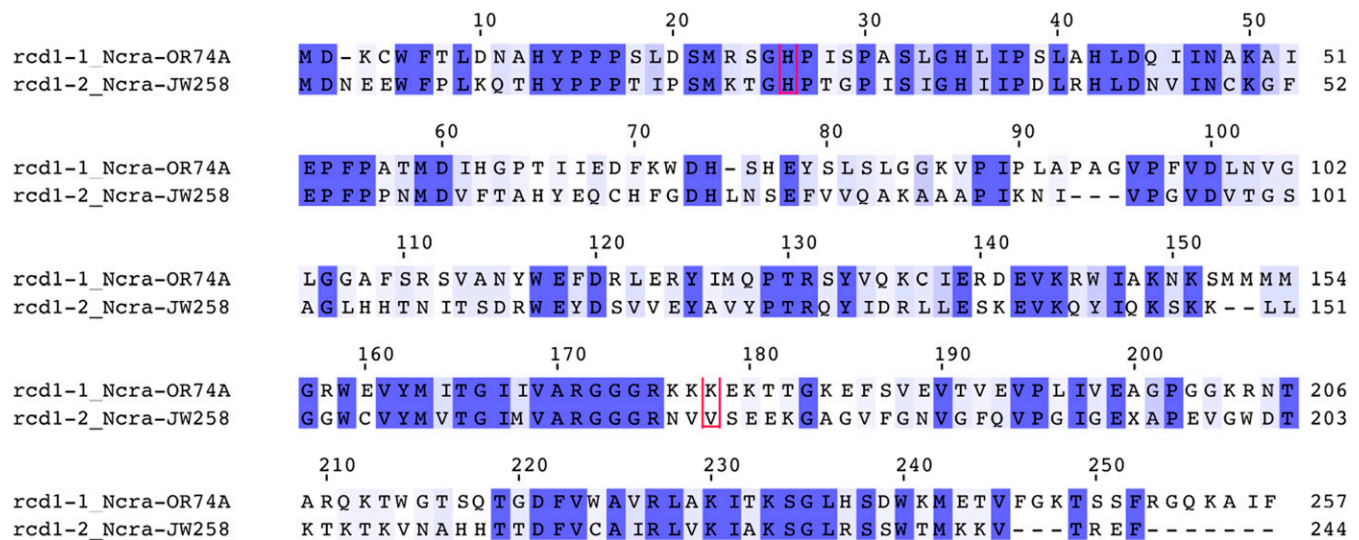
The similar GRD phenotype between the two loci that regulate death upon fusion of germlings, *sec-9/plp-1* and *rcd-1*, suggested that a functional relationship might exist in the induction of GRD by allelic differences at these loci. Therefore, we assessed whether the GRD phenotype induced in germlings with different *sec-9/plp-1* allelic specificities was dependent on *rcd-1*, by assessing *sec-9/plp-1*-induced GRD in a  $\Delta$ *rcd-1* mutant. Genetic differences at *sec-9/plp-1* were sufficient to induce GRD in a  $\Delta$ *rcd-1* mutant background, with a frequency of cell death and cellular phenotype that was



**Figure 2** NCU05712 (*rcd-1*) is necessary and sufficient for GRD induction in *N. crassa*. (A) A flow cytometry-based quantification of GRD using the vital dye PI (Heller *et al.* 2018) shows that *rcd-1* is necessary for the GRD phenotype. (B) GRD quantification with isogenic strains of *N. crassa* carrying different *rcd-1* alleles. Strains carrying ectopic *rcd-1* allele at the *his-3* locus are indicated with (h::) preceding the allele. Experiments were performed in triplicate, with 20,000 events counted per experiment. \* $P < 0.0006$ , one-way ANOVA with Tukey's multiple comparisons test. (C) Reconstitution of *rcd-1-1/rcd-1-2* GRD phenotype in isogenic *N. crassa* strains. Cells undergoing GRD (white arrows) show strong vacuolization and uptake of PI. Deletion of *rcd-1* abolishes GRD and genetically distinct strains can undergo cell fusion to form viable cell networks. Fusion points between genetically distinct germlings are shown with black asterisks. Bar, 5  $\mu$ m.

indistinguishable from strains inducing *sec-9/plp-1*-triggered GRD in a wild-type genetic background (Figure S5). Thus, RCD-1 does not function downstream of SEC-9/PLP-1 in the execution of cell death, suggesting that these two GRD pathways act independently.





**Figure 3** Alignment of GRD-inducing allelic variants RCD-1-1 and RCD-1-2 from the wild-type reference strain FGSC2489 and the wild isolate JW258. Identity between the two proteins is shown in dark blue, similarity in light blue. Amino acids residues found to be under positive selection are boxed in red.

### Genomic rearrangements at the *rcd-1* locus in *sordariaceae*

The *rcd-1-1* and *rcd-1-2* alleles encode proteins of unknown function of 257 and 244 amino acids, respectively. The two allelic variants showed 38% identity (54% similarity) on the protein level (Figure 3). The coding region of both *rcd-1* alleles consisted of two exons, as confirmed by sequencing *rcd-1-1* cDNA from FGSC2489. The analysis of *rcd-1* gene structure showed a partial duplication of *rcd-1-1* in the 3' end of the *rcd-1-2* allele in the JW258 strain. The duplicated *rcd-1-1* in the *rcd-1-2* background lacked 212 bp from the 5' end of the ORF, including the start codon of the gene and the promoter (Figure 4). We examined the *rcd-1* locus of other *rcd-1-2*-encoding *N. crassa* strains, and found that all sequenced wild isolates of the *rcd-1-2* genotype carried a partial duplication of the *rcd-1-1* allele at the 3' end of *rcd-1-2*. However, unlike JW258, the other *N. crassa* *rcd-1-2* strains have a shorter partial duplication of the *rcd-1-1* allele (*rcd-1-1*<sup>Δ1-610</sup>), comprising mostly the second exon (Figure 4). These data suggest that an *rcd-1-2* allele inserted into an *rcd-1-1* allele, thereby disrupting the gene structure of the latter.

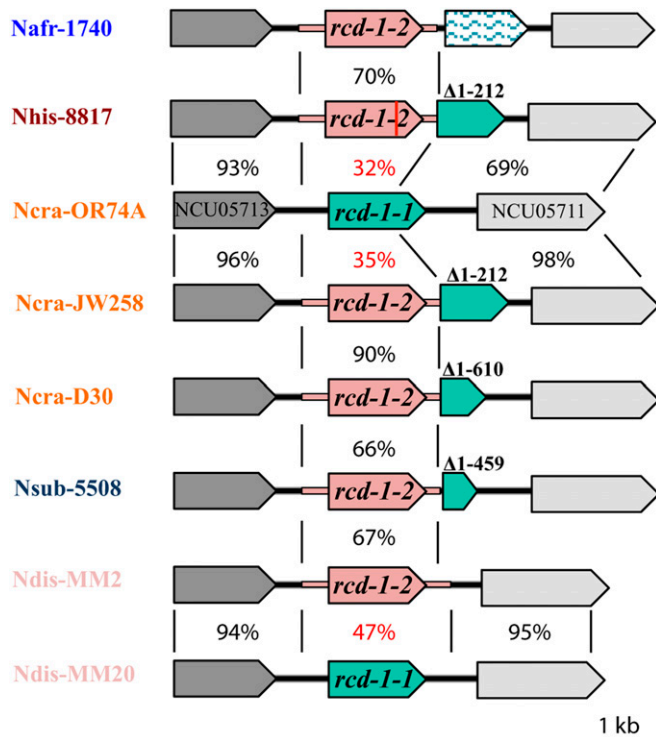
Sequenced *N. crassa* strains carrying the *rcd-1-2* allele had a partially duplicated *rcd-1-1*<sup>Δ1-610</sup> or *rcd-1-1*<sup>Δ1-212</sup> copy, with the vast majority of strains carrying *rcd-1-1*<sup>Δ1-610</sup> and only JW258 presenting the longer duplication (Figure S6). Yet, the insertion site in the intergenic region upstream of the *rcd-1-2* locus in sequenced *rcd-1-2* *N. crassa* strains was the same, strongly suggesting that one ancestral insertion event was the origin of the *rcd-1-1* gene disruption in *N. crassa* and that the partially duplicated *rcd-1-1*<sup>Δ1-610</sup> and *rcd-1-1*<sup>Δ1-212</sup> alleles evolved during a subsequent independent event (Figure S7). The molecular signature of the *rcd-1* gene disruption event was also found in the genomes of other related *Neurospora* species. For example, in the genome of *N. sublineolata*

(FGSC5508), downstream of the *rcd-1-2* allele was a partially duplicated *rcd-1-1*<sup>Δ1-459</sup> sequence, while in *N. hispaniola* (FGSC8817), the *rcd-1-2* allele had the same insertion sites as in *N. crassa* JW258 strain and thus carried a partially duplicated *rcd-1-1*<sup>Δ1-212</sup> allele (Figure 4 and Figure S6). This observation suggested that the genomic rearrangement was a relatively ancient event, and likely predates speciation. Although, we did not find a partially duplicated *rcd-1-1* ORF in *N. discreta* *rcd-1-2* strains, we identified clear breakpoints in the sequence homology of the *rcd-1* locus between *N. discreta* strains carrying *rcd-1-1* and *rcd-1-2* alleles (Figure S8). As the sudden decrease in sequence similarity was in the upstream/downstream region of the *rcd-1-2* gene, similarly to the *N. crassa* strains, we concluded that, in *N. discreta*, the insertion of *rcd-1-2* led to the complete loss of the *rcd-1-1* ORF. Likely, taking into account the observed degeneration of the pseudogenic partial *rcd-1-1* alleles in the *rcd-1-2* strains of *N. crassa* and *N. hispaniola*, the *rcd-1-2* locus in *N. discreta* strains has resulted from the same initial ancestral event of an *rcd-1-2* allele disrupting an *rcd-1-1* locus. These data suggest that the partial *rcd-1-1* allele present in the *rcd-1-2* strain was not sufficient to induce GRD. Consistent with this hypothesis, we quantified the amount of cell death in mixtures of germlings expressing the *rcd-1-1* allele, and germlings expressing the *rcd-1-2* allele lacking the partial duplication of *rcd-1-1* (*rcd-1-2-NPD*) or *rcd-1-2* in the presence of partially duplicated *rcd-1-1*<sup>Δ1-610</sup>. A significant difference in the frequency of cell death between these two germlings mixtures was not observed (Figure S9).

### Evolutionary signatures of balanced and long-term positive selection associated with *rcd-1*

As *rcd-1* is a novel allorecognition determinant, we investigated the evolutionary signatures associated with the gene.





**Figure 4** Genetic rearrangements in the *rcd-1* locus in *Neurospora*. *rcd-1-1* (green) and *rcd-1-2* (pink) alleles from different species are shown as cartoons in the *rcd-1* locus. The extended region in pink in *rcd-1-2* represents the sequence with sharp decrease in homology (or regions with lack of homology) at nucleotide level between strains of the *rcd-1-1* and *rcd-1-2* genotype. Percentage identities between two different genomic regions are shown between black lines delimiting the comparisons. Numbers over the *rcd-1-1* allele indicate the limits of the truncated ORF in the genome of different *Neurospora* species and strains. Red line in the *rcd-1-2* allele of *Neurospora hispaniola* (Nhis-8817) represents a premature stop codon. Species names are abbreviated as follows; *N. africana* (Nafr), *N. hispaniola* (Nhis), *N. crassa* (Ncra), *N. sublineolata* (Nsub), *N. discreta* (Ndis).

Natural selection has been shown to operate on allorecognition loci via a mechanism of negative frequency dependent selection that favors rare alleles and results in the maintenance of multiple alleles at relatively high frequencies (Wu *et al.* 1998; Hall *et al.* 2010; Zhao *et al.* 2015). The high positive Tajima's D scores calculated for *rcd-1* in *N. crassa* ( $D = 3.491$ ) and *N. discreta* ( $D = 2.637$ ) indicated an excess of intermediate frequency alleles at *rcd-1* in the two *Neurospora* populations (Table 1). To deepen our analysis of the allelic distribution of *rcd-1*, we investigated the phylogenetic history of this locus in members of the Sordariaceae. We analyzed 194 fungal genomes from 26 species and identified 113 *rcd-1* homologs in 104 of the sequenced strains (Table S1). Two Chaetomiaceae species (*Chaetomium cochliodes* and *Thielavia hyrcaniae*) and one Lasiosphaeriaceae II had two copies of *rcd-1*, and one Lasiosphaeriaceae IV (*Cladorrhinum bulbillosum*) had seven copies. However, *rcd-1* homologs were not identified in 90 genomes, representing 12 species of Chaetomiaceae and Lasiosphaeriaceae I and II. We found that 81 of these genomes belonged to

*N. tetrasperma* (65) and *N. discreta* (16) strains, where gene models encoding longer versions ( $>500$  aa) of *rcd-1* were predicted in different lineages, likely corresponding to annotation errors. When examined manually, *rcd-1* alleles from *N. tetrasperma* strains (FGSC7586, FGSC9035) carried premature stop codons, similar to an *rcd-1* allele from *N. hispaniola* (FGSC8817).

Phylogenetic analysis (maximum-likelihood) of *rcd-1* in 27 wild isolates of *N. crassa* from Louisiana revealed a nearly equal distribution of two main polyphyletic clades (14 *rcd-1-1*-like and 13 *rcd-1-2*-like homologs in each clade) (Figure 5A and Table S1). Furthermore, in *N. discreta*, a relatively equal distribution (60–40%) between the two clades for the 35 analyzed *rcd-1* homologs was also identified. The lack of monophyletic segregation between the *N. crassa* and *N. discreta* alleles indicates that the allele-specific polymorphisms predate the speciation event, and represent an example of *trans*-species polymorphism (Figure 5). Other *Neurospora* species carrying *rcd-1* alleles also split between the two clades with *N. pannonica* and *N. sitophila* *rcd-1* alleles being closer to *rcd-1-1* alleles, while *N. terricola*, *N. africana*, and *N. sublineolata* carried sequences situated on the *rcd-1-2* branch. The allele-specific polymorphisms were distributed over the entire length of the allelic variants (Figure S10). We did not find marks of *trans*-species polymorphism on the adjacent genes (NCU05713 and NCU05711) to *rcd-1* (NCU05712), which are situated at  $\sim 1.3$  kb and  $\sim 0.9$  kb, respectively, from NCU05712 (Figure S11). The equal distribution of allelic frequencies in the wild populations of *Neurospora* and the observed *trans*-species polymorphism suggest that the *rcd-1* locus is under long-term balancing selection.

Positive diversifying selection has been reported previously to operate on different allorecognition genes in *Neurospora* and *Podospira* (Bastiaans *et al.* 2014; Zhao *et al.* 2015). Sites under positive selection could be of importance for subsequent functional analyses of the *rcd-1* allorecognition process. We thus tested for long-term positive selection by comparing, with likelihood-ratio tests, models assuming positive selection with null models in PAMs codeml program. For site-specific analyses, positive selection model M8 (which assumes that one class of sites has  $\omega > 1$ ) provided a better fit to the data than purifying selection models M7 and M8a (which assumes  $0 < \omega < 1$  and  $0 < \omega \leq 1$ , respectively; likelihood ratio test:  $P = 0.0397$  for M7 vs. M8,  $P = 0.0374$  for M8a vs. M8) (Table 2). Under M8a, a single amino acid had a  $>0.95$  posterior Bayesian probability of being under positive selection using the Bayes empirical Bayes method. For branch-site analyses, positive selection model A (which assumes positive selection  $\omega > 1$  along foreground branch *rcd-1-2*) provided a better fit to the data than purifying selection model A (which assumes  $0 < \omega \leq 1$  along all branches; likelihood ratio test:  $P = 0.0003$ ). Under model M8 and A, respectively, one and two amino acids (plus the stop codon) had a  $>0.95$  posterior Bayesian probability of being under positive selection using the empirical Bayes method. Both models found the position 173 (V173 in RCD-1-2; boxed in



**Table 2** Parameter estimates and likelihood scores of negative selection (A1, M7, and M8a) and positive selection models (A, M8)

Model	dN/dS	Parameter estimates	PSS	Likelihood
M7	0.364	$p = 1.06499, q = 1.81851$	NA	−7758.9
M8	0.365	$p0 = 0.99506 P = 1.13305 q = 2.02613 (p1 = 0.00494) \omega = 2.65855$	1 (1)	−7755.6
M8a	0.359	$p0 = 0.96236 P = 1.21055 q = 2.35599 (p1 = 0.03764) \omega = 1.00000$	NA	−7757.8
A1		Site class 0 1 2a 2b Proportion 0.75587 0.20934 0.02724 0.00754 Background w 0.24450 1.00000 0.24450 1.00000 Foreground w 0.24450 1.00000 1.00000 1.00000	NA	−7774.8
A		Site class 0 1 2a 2b Proportion 0.76963 0.21413 0.01270 0.00353 Background w 0.25112 1.00000 0.25112 1.00000 Foreground w 0.25112 1.00000 15.33251 15.33251	4 (3)	−7764.5

The dN/dS ratio is an average across all sites categories. PSS, number of positively selected sites (number of sites with posterior probability cutoff  $P > 95\%$ ).

genes. These results suggest that the *rcd-1* allorecognition process operates in the wild.

### Distribution of *rcd-1* homologs in Ascomycota

Functional allorecognition genes are generally poorly conserved between different fungal species (Fedorova *et al.* 2005; Paoletti 2016; Daskalov *et al.* 2017). However, genes encoding proteins carrying predicted domains associated with allorecognition processes fall in several large gene families and are distributed throughout fungi (Van der Nest *et al.* 2014; Zhao *et al.* 2015; Paoletti 2016). As *rcd-1* is not related to any currently known fungal cell death-inducing determinant, we decided to explore the broader distribution of *rcd-1* homologs. BLAST searches with either of the allelic variants of RCD-1 retrieved ~260 significant hits (E value <0.05) from the nonredundant proteins database (NCBI). All identified sequences were encoded in the genomes of filamentous ascomycetes (Pezizomycotina), predominantly in species in the Sordariomycetes. The number of *rcd-1* homologs increased to ~940 sequences when we performed a profile HMM search using HMMER with RCD-1-1 as initial query sequence (Finn *et al.* 2011; Potter *et al.* 2018); the HMM search was performed until no new sequences were retrieved, which occurred after the third iteration. Almost all significant hits (>99.9% of hits with E value <0.005) were distributed in 24 fungal orders (23 in Ascomycota and 1 in Basidiomycota) and three bacterial orders containing only four sequences (Figure 6 and Table S2). The unique *rcd-1* homolog found in Basidiomycota was from *Naematelia encephala* (Tremellales), while the prokaryotic sequences were distributed in three Gram-negative species (Flavobacteriales and Xanthomonadales) and one *rcd-1* homolog in the filamentous actinomycete *Actinoplanes missouriensis* (Micromonosporales) (Table S2). The four bacterial sequences grouped near the root of the tree in two different nodes with other fungal *rcd-1* homologs (Figure 6). Two fungal orders—Hypocreales (324) and Eurotiales (112)—contained approximately half of the analyzed *rcd-1* homologs. Yet, *rcd-1* genealogy did not follow Ascomycota phylogeny, because a large number of nodes grouped *rcd-1* homologs from phylogenetically

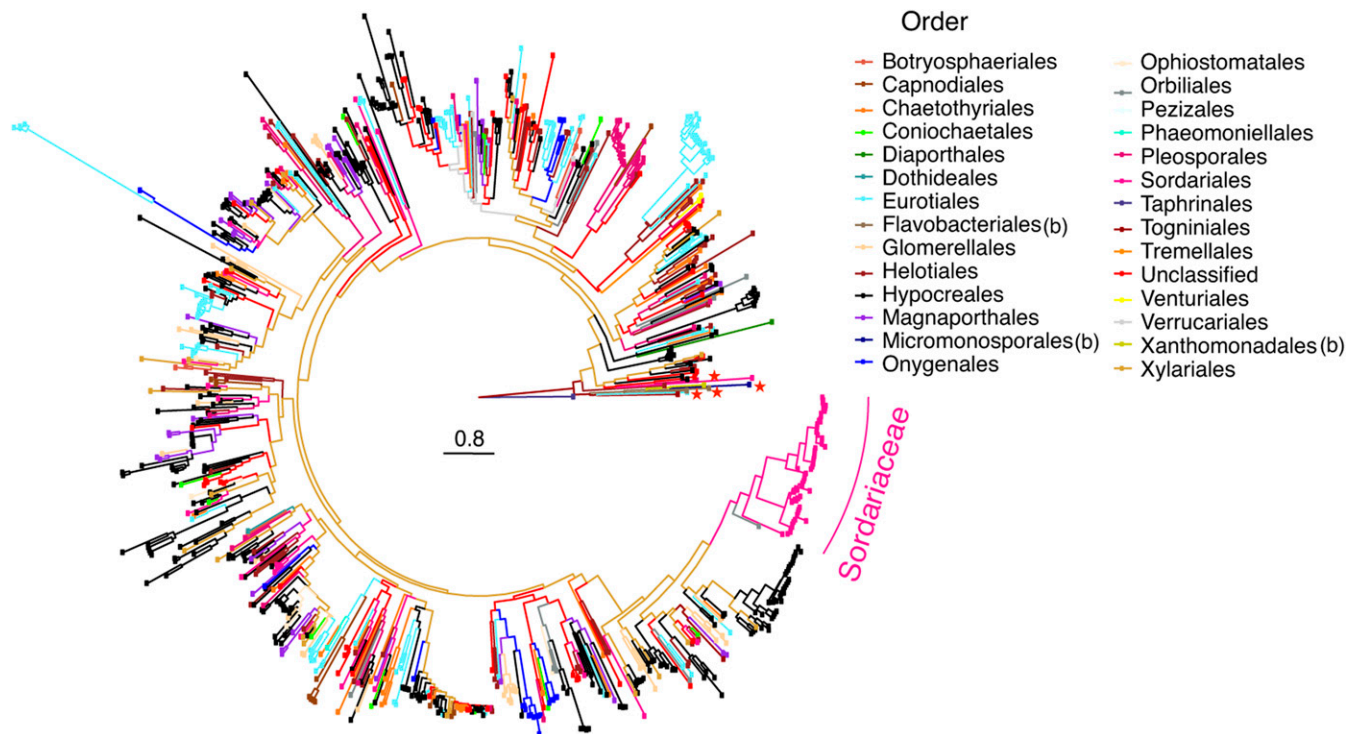
distant taxa (Figure 6) [https://figshare.com/articles/Programmed\\_Cell\\_Death\\_in\\_Neurospora\\_crassa\\_Is\\_Controlled\\_by\\_the\\_Allorecognition\\_Determinant\\_rcd-1/10298045](https://figshare.com/articles/Programmed_Cell_Death_in_Neurospora_crassa_Is_Controlled_by_the_Allorecognition_Determinant_rcd-1/10298045). For example, *rcd-1* homologs from Hypocreales (black in Figure 6) are interspersed among homologs from other orders of Ascomycota and are found on most branches of the tree. A similar distribution is observed for homologs belonging to Eurotiales (sky blue), Magnaporthales (purple), and, in general, most orders with high number of *rcd-1* homologs (Figure 6). Sequences from Sordariales (pink) were scattered in distant nodes of the tree, with previously analyzed *rcd-1* homologs from the Sordariaceae (mainly *Neurospora* species) (Figure 5A)—where the incompatible *rcd-1-1* and *rcd-1-2* alleles could be found—grouped together in a separate branch of the tree. The latter finding was not surprising as the sequences situated on the Sordariaceae branch are alleles of *rcd-1* from one species (*N. crassa*, *N. discreta*, and *N. tetrasperma*) and/or *rcd-1* orthologs from closely related species.

At least one *rcd-1* homolog was present in ~180 fungal species, with a mean of five and median of three *rcd-1* homologs per strains (sequenced genome). The genomes of more than a dozen ascomycete taxa contained >10 *rcd-1* homologs. Among the species with higher numbers of *rcd-1* homologs were several *Trichoderma* species, where individual strains carried up to 23 *rcd-1* genes in their genome (*T. atroviride* ATCC 20476). Remarkably, the genome of the closely related species *T. reesei* QM6A contained only two *rcd-1* genes. We concluded from these results that the allorecognition determinant *rcd-1* identified in *N. crassa* is a member of a large gene family, widespread in the Ascomycota. The phylogeny of *rcd-1* in fungi suggests that the gene has experienced lineage-specific gene expansions and was subject to horizontal transfers between species. Similar distribution patterns have also been reported for other families of nonself recognition determinants in fungi (Dyrka *et al.* 2014; Zhao *et al.* 2015).

### Discussion

Here, we investigated the molecular basis of programmed cell death occurring during nonself recognition in germinating





**Figure 6** Phylogenetic distribution of *rcd-1* in fungi and bacteria. Maximum likelihood tree of ~940 *rcd-1* homologs. Branches of the phylogenetic tree are color-coded in accordance with the taxonomic rank at order level for the species from which the various *rcd-1* sequences have been retrieved. Order names are indicated in the panel next to the phylogenetic tree. The gene family expansion in Sordariaceae family (order Sordariales), which display the various *rcd-1-1* and *rcd-1-2* *Neurospora* alleles from this study, is highlighted. Branches corresponding to the four bacterial sequences are indicated with red stars. Bacterial phylogenetic orders are indicated with the letter (b) in the legend of the figure. Sequences from species with no clearly established phylogeny at order level, are grouped under “unclassified”.

conidia of *N. crassa*, and identified NCU05712 as a novel allorecognition determinant, which we named *regulator of cell death-1* or *rcd-1*. Two incompatible alleles, *rcd-1-1* and *rcd-1-2*, were identified at the *rcd-1* locus and the coexpression of these alleles in fused germlings and hyphae was necessary and sufficient to trigger massive vacuolization and cell death. The two alleles showed evidence of balancing-selection and were associated with *trans*-species polymorphism, two molecular hallmarks associated with genes involved in the control of nonself discrimination (Richman 2000; Hall *et al.* 2010; Milgroom *et al.* 2018). Furthermore, close examination of the *rcd-1* locus revealed that an *rcd-1-2* allele inserted in an *rcd-1-1* allele, disrupting the gene structure of the later. We find this observation particularly intriguing as it underscores a trend of genomic rearrangements frequently associated with nonself recognition loci. For example, several distinct GRD-specific haplotypes are present at the recently identified *plp-1* locus in wild populations of *Neurospora* and *Podospira* (Heller *et al.* 2018). Multiple rearranged haplotypes were also found at the *doc-1* (*determinant of communication-1*) locus, which defines nonself recognition at distance (precontact) in germlings (Heller *et al.* 2016), while, in *Cryptonectria parasitica*, the incompatibility locus *vic4* contains two idiomorphic genes (Choi *et al.* 2012). Population genomics data and evolutionary signatures of *rcd-1* suggest

that the genomic rearrangements may have initiated events that led to the emergence of this incompatibility system in at least some *Neurospora* species. This hypothesis is based on the following observations: (i) *rcd-1* alleles appear to be under balancing selection, evenly distributed in the populations of *N. crassa* and *N. discreta*, which suggests that natural selection is acting on the *rcd-1-1/rcd-1-2* incompatibility; and (ii) all currently sequenced *rcd-1-2* strains from *N. crassa* and *N. discreta* carry molecular evidence of a rearrangement in the *rcd-1* locus. If the *rcd-1-1/rcd-1-2* incompatibility system predates the rearrangements in the *rcd-1* locus, and, assuming that the adaptive value of a hypothetical wild-type *rcd-1-2* and the “inserted *rcd-1-2*” is equivalent, one could ask why the locus carrying the insertion is so strongly over-represented in *N. crassa* and *N. discreta*. A plausible explanation is that the *rcd-1-2* allele has been horizontally transferred or introgressed; this event could have been possible only if it disrupted the *rcd-1-1* allele (coexpression of both alleles is lethal).

Interspecies transfer of HI genes and mating-related genes has been previously documented in *Ophiostoma novo-ulmi* (Paoletti *et al.* 2006). In *Neurospora*, introgression has been shown to play a role in shaping mating-type chromosomes, and, possibly, the evolution of the *rsk* (resistant to spore killer) locus (Sun *et al.* 2012; Svedberg *et al.* 2018). Further



phylogenetic analyses are needed to explore the evolutionary origin of *rcd-1* incompatibility, notably sequencing of additional *Neurospora* strains and/or species closely related to *Neurospora* within the Sordariales. The accumulation of such genomic data could identify more *rcd-1* haplotypes from species with population samples, and offer a better view of the allelic diversity of *rcd-1* in Sordariales, which could reveal a more precise hypothesis regarding the emergence of *rcd-1-1/rcd-1-2* incompatibility. The discovery of the *rcd-1* incompatibility system represents an exciting opportunity to investigate the evolution of allorecognition genes and how particular genomic rearrangements impact their distribution in fungi.

Several key contributions have been made recently toward a better understanding of the molecular basis of allorecognition in *N. crassa*, with three key steps or checkpoints that regulate early colony establishment (Dyer 2019). The first checkpoint regulates chemotropic interactions between germlings, with *Neurospora* populations showing five communication groups, and which is regulated by allelic specificity at the *determinate of communication* or *doc* loci. Isolates from the same communication group (and with identical *doc* allelic specificity) show robust chemotropic interactions, while isolates from different communication groups (with different *doc* allelic specificity) show significantly reduced communication, and, thus, cell fusion (Heller *et al.* 2016). The second checkpoint, controlled by the *cwr-1* and *cwr-2* (cell wall remodeling) loci, regulates whether cell wall dissolution and cytoplasmic mixing occurs between adhered germlings (Gonçalves *et al.* 2019). Cells with identical allelic specificity at *cwr-1* and *cwr-2* undergo cell fusion and cytoplasmic mixing, while adhered germlings with alternate *cwr-1 cwr-2* allelic specificity are blocked. If cells have identity at the *doc* and *cwr* loci, somatic cell fusion proceeds, but the viability and fitness of fused germling are dependent upon the GRD loci – *plp-1/sec-9* (Heller *et al.* 2018) and *rcd-1*. These loci also control allorecognition in mature hyphae and thus could play a critical role in fungal interindividual relations beyond germling interactions and colony establishment.

In future work, investigations into the molecular mechanisms of *rcd-1*-induced cell death are of major interest as the gene is active in a developmental stage (asexual spores) frequently associated with threats to human and plant health. Targeting of endogenous cell-death pathways in fungi could represent a viable strategy in the fight against various fungal pathogens; it was recently proposed that mice clear inhaled spores from their lungs by targeting fungal PCD (Shlezinger *et al.* 2017; Kulkarni *et al.* 2019). The identification of *rcd-1* diversifies the inventory of allorecognition genes identified in fungi. Considering the widespread nature and abundance of *rcd-1* homologs, especially in the Pezizomycotina, we hypothesize that conspecific nonself discrimination is unlikely to be the sole role performed by *rcd-1* in all of these species. This hypothesis is especially intriguing in light of recent reports positioning various allorecognition determinants inside broader gene families, which control innate immunity in

animals and plants (Paoletti and Saupe 2009; Dyrka *et al.* 2014; Gonçalves *et al.* 2017). Future work on the molecular mechanism of recognition and induction of cell death mediated by RCD-1 will test its role in the paradigm of allorecognition and potentially provide clues to alternative functions of this gene family in the Pezizomycotina.

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